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Effect of tRNA Availability on the Rate of Elongation of Translation is a Non-uniform Process Nascent Polypeptide Chains STANISLAS VARENKE, JEAR BUC, ROLAND LLOUBES AND CLAUDE LAZDUNSKI Centre de Biochimie et de Biologie Moléculaire du C.N.R.S. B.P. 71, 13402 Marseille Cedex 9, France

(Received 2 December 1983, and in revised form 18 July 1984)

We reported elsewhere (Varenne et od., 1982) that, during synthesis of a number of in Bacherichin cali, intermediate naucent chains of discrete sixes accumulated, suggesting a variable rate of translation. In this paper, a detailed analy as provides arguments that this phenomenon, at least for the proteins under study, is not related to aspects of messenger RNA such as secondary arructure. It is linked to the difference in transfer RNA availability for the various codons. Experimental analysis of translation of other proteins in E. ooli cunfirms that the main origin for the discontinuous translation in the polypeptide elongation cycle is the following. For a given codon, the stochastic search of the cognate ternary (aminoneri-tRNA-EF-Tu-GTP) is the rate-limiting step in the (RNA concentrations. The resistention of this model and its possible clongation cycle: transpeptidation and translocation aters are much degree of slackening in ribosome movement is almost physiological significance are discussed romplex

1. Introduction

The idea that intracellular concentrations of tRNAs play an important role in the Ames & Hartman (1903). Anderson (1969) proposed that the rate of translation tRNA species, energies of interaction between codons and anticodons, has led to a kemura, 1981a,5,1982; Grosjean & Ficra, 1982). The idea that translation occurs dynamics and the regulation of protein synthesis was suggested 20 years ago by might be slowed in vito at the site of regulatory codons. Since then, more detailed knowledge of nucleatide sequences, tRNA concentrations, decoding spectra of number of authors emphasizing the importance of IRNA concentrations and/or of codon-anticodon interaction energies in the dynamics of translation le.g. 2008 Grantham et al., 1981; Chavancy & Garel, 1981; Gouy & Cautier, 1982. at a variable rate is implicit or explicit in their work, but to our knowledge, no study has yet proposed a quantitative relationship between tRNA concentration and rate of elongation at each codon.

it has been observed in our laboratory that elongation of nascent polypeptide hains of colicin A, El, E2 and E3 occurs at a variable rate (Varenne et al., 1982;

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established (Morlon et al., 1983), and since ecdicin A is highly expressed after induction, whereas synthesis of chromosomal proteins is strongly reduced, colicin A synthesis constitutes a good model system for an experimental approach to the azdunski et et., 1984). Since the nucleotide sequence for colicin A has been dynamics of the clongation eyele.

In this work, we have analysed in detail the effect of tRNA availability on the inhypeptide elongation rate for coliein A and for some other Bischerichia coli incteins. Our results demonstrate directly two important points: (1) the rate-limiting tRNAs exist in E. coli in different intracellular concentrations (even those preferentially used by highly expressed proteins), the rate of translation varies along the mRNA Moreover, for a given protein, complete translation of step in the elongation each of polypeptide is tRNA selection; (2) as the different individual mRNAs occurs at different rates.

The possible effect of codon-anticodon energies of interaction has not bren taken into account for lack of adequate data. However, our results indicate that the role of this factor in the rate of translation, if it existed, would have to be less insportant than that played by tRNA concentrations.

2. Materials and Methods

(a) Materials, bacterial strains, grouth conditions and conditions of radio la bellina

exactly as described (Carenne et al., 1982). For praduction of colicin E1, the strain E. coli All materials were as described (Varenne et al., 1982). Procedures were also carried out KI2 W3110 Col El. 100f was used.

(b) Preparation of somples for impreparecipitation

In addition to the previously reported experimental protocol, another technique was occasionally used to solabilize cell pellets. These were taken up in 10 pl of 2%, (w/r) sodium dodectd sulphate and incubated for 5 min at 180°C, allowed to cool shouly and 50 pl of the irammoprecipitation buffer (without sodium dedexy sulphase) added us described Varenne (1 al., 1982)

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except for a pre-exposure of the film to a hypersensitizing light flash resulting in an increase of background film absorbance of about 0.13 at 510 nm. This treatment permits quantitative interpretation of film density (Laskey & Mills, 1975), using, if necessary, electrophoresis and fluorography were also carried out as described (Farenne et al., 1982), antisers were ultained as described (Varenne et al., different exposures for very contrasted fluorograms. Antibudies and

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the influence of tRNA availability on the dynamics of protein synthesis. The clungation , Gour & Grantham (1980). Chavaner & Garel (1981) and Gony (1981) have pointed out excle is described by (sour & Gantier (1982) thus:

In and GTP) diffusing in the evitablism interact with the codon and the ribosome at the therefore the aminoacyteRNA dissociates from the ribusome. When the specificity Thus. each codon can be characterized by the average number of codon-tRNA interactions at the A-site during one clongation cycle. The teletive concentration of the endon-cognate tRNA is equivalent to its probability of colliding with the Asite codon. Hence, if this probability is C the mean number of ordon-tRNA interactions necessary for the elongation ibosomal A-site. The tencary complexes (ammoney) tRNAs bound with elongation factor A-site. Most often the codon does not belong to the IRNA recognition spectrum and condition is fulfilled, the elongation ercle starts: transpeptidation and translocation occur. is found at the the beginning of each polypeptide changation eyele, a codon eyele to occur is L/C, "

regnate tRNA from the A-site and the next collision with a tRNA. Then the mean duration of the addition of a given araino acid residue corresponding to a given codon is $I=(\theta_0+\theta_1)\lambda^3+I_2+I_3$, θ_1 is independent of the codon considered. If we assume that θ_0,I_2 be the mean direction of an interaction between a given codon and a non-cognate tRNA at The relationship between f, the mean duration for the addition of a given amino seirl residue corresponding to a given codon, and I/C (I/C will be called N) is the following: I is the sum of 3 mean durations: I, for the search of the adapted terms γ complex for the codon in the A-site of the ribosome; t_2 for transpeptidation; and t_3 for translocation. Let $heta_6$ the A-site of the ribosume, and let $heta_i$ be the mean duration between the ejection of a nonand to ave also independent of the codon, then t=3.N+B, where A and B are constants.

(e) Determination of this 4 concentrations and for computation of N

corresponding codons is more or less strong according to whether the particular protein is highly or weekly expressed. Brating this in mind. Gouy (1981) has established an average tRNA usage by considering that (10%, of total proteins can be considered as highly Since concentrations of only nome tRNA species have been determined experimentally, it nas preessary to calculate an approximate value for the remainder. We have taken advantage of the fact that a good correlation exists between the abundance of tRNAs and their usage in the cell (Ikemana, 1981a.b. Gony, 1981). It is thus possible to determine by regression jakt of experimentally determined IRNA roncentrations as a function of usage of these LRNAs in total E. role proteins. This general usage can be approximated from the garticular usage for a number of inPNAs for which sequences are known. The degree of correlation between the abundance of IRNA species and the usage of the supposedly interjedation an approximate value of unknown IRNA concentrations by using the linear expressed and 40% as weakly expressed.

overriding importance since, in each class, each protein uses IRNAs in approximately the same way (Ikemura, 1981a.b). On the same granuds, we have exhablished a new encloss we have used 21 sequences, or part sequences, corresponding to highly expressed genes (Gany & Gautier, 1982), and 41 sequences of weekly expressed genes. The cordon usage for the latter comes from the Livon sequence bank ACNUC. Among these 41 genes, 28 are In each class, each protein is involved only by the number of times that each of the fil exclores appears in its sequence and not by its nammit in the cell. However, this is not of usage [Table I], which differs slightly from that established by Gony (1981) for 2 massum [1] extrachromosomal genes were not taken into account, since takes genes are not expressed permanently in E. coli: (2) aumerous new sequences have been determined and referred to by Gouy & Gautier (1982). They are: nust., Incl., Incl., ifiG. 1904. 190B. 1901). trpd. trpl3, trpG, trpD, trpE, trpR, thrd, and1, une2, ane3, ane4, ane6, anec, anec. nuec. aneli. uneG, aneli. goff (Von Wikken-Bergmann & Maller-Hill, 1982), Inwis (Chinent & Hofmung, 1981), JudA 1981), the Kossari & Giegnal-Sanzay, 1982; Ailia et al., 1982), fel (Smith & Calvo, 1980), thank (Deeley & Yannishky, 1981), typs (Hall et al., 1982), 13 Kit and 15 Kit princins linyte et al., 1982), rpol. (Oxelimnikov et al., 1982), thrB and thal (Cassart et al., 1981)

4RNAs, the repartition between these 2 1RNAs has been carried out as indicated by Ikemura (1981a). The correlation between the amounts of 1RNAs and their frequency of by interpolation from the regression line corresponding to the plot of amount of tRNA corresponding to the plot of amount of tRNA corresponding obtained and used in concentrations, the total is equal to 1. The 6 concentrations concerning codons UCU (Ser), CUC (Cir.), ACU (Thr), CCU (Pro), GCU (Ala) and CUU (Val) are apperent concentrations endon usinge corresponding to the 62 genes has been allocated between tRNAs hy ussee for 23 (RNAs quantified by Ikernura (1981a) is shown in Fig. 3. The correlation coefficient of 0.86 indicates that one can coughly estimate the other IRNA concentrations concentrations (experimentally determined or interpolated). Including the tRNA,441 using the decoding spectrum proposed by Exercina (1981a). For codous recognized by 2 isofurther calculations are indicated in Table 2. Thirty-five concentrations are actual that take into secount the recognition of these cusions by 2 different IRNAs.

In a number of rainulations, the value for the tRNAmen has been replaced by the

interpolated value. The latter was obtained from the regression line derived from only 22 points, but was very close to that obtained with 23 points.
Grosjean & Piers (1982) proposed a decoding spectrum that differs from that of Ikemura. (1981a) for 6 codois; (1) seconding to these authors, 2 different tRNA Leu iso-acceptors translate codons UUA and UUG: as the codon usages are quite similar (about 50% versus allo,), the experimentally determined concentration has been distributed equally between these 2 tRNAs: (2) according to Groupsan & Fiera (1982), it is not certain that codons AAG (Lys) and GAG [Glu] can be decoded by the tRNAs decoding AAA [Lys] and GAA (Glu), respectively. In this hypothesis, we have assumed that experimentally determined concentrations concerned the sum of the 2 tRNAs, and these concentrations have been distributed between these 2 tRNAs proportionally to the codon usage. In the hypothesis where the experimentally determined value was related to the major tRNA, the conventrations should be modified acouchingly, but this has minor consequences for futther

ridiscone. I wise abring one elongation cycle sabbreeisted to N) as a function of the migration (1) Onta treatment: determination of the neeringe animher of codon-IRNs inhermetions at the of the curresponding elongation intermediates.

Basic programs were developed with a Wang 2000 microcompater provided with a thippy thisk and a digital plotter.

As a first step, fluorograms obtained (10 cm migmition) were enlarged a times unider conditions that preserve contrast. The enlargement was acanned with a 4 times expansion in migration. Knowing the migration of calibration standards, one can deternine coefficients a and b for the migration according to $x\approx a\log M_r+b$.

diffusion of polypeptides in the gel, light-scattering in the fluoringram and stit width), it is necessary to simulate this diffusion. This dispersion can be accomplished through a for cuton i, routinely called concentration for convenience). After computation, the plot zq (abscissa). N fordinate) is drawn. As a certain dispersion in densitometer profiles exists For each codon (C.) the computer determines the cumulated weight of all amino acid residues assembled until the corresponding amino acid (A), the corresponding position in the scanning (z,) and the number N_i (1/ G_i , where G_i is the frequency of the tRNA specific Gaussian distribution with a full width at half maximum (FIPHM) either constant in distance ($\Delta r = c(e)$, or constant in M, ($\Delta M_e = c(e)$, A convolution (called dispersion in the lext and denoted Na) from the direct calculations is thus obtained.

Another computer program plots x_i (abserve). N_i (ordinate) with $z_i = C(i+d)$, and dispenses the results with a full width at half maximum constant in amino acid residue number (di = 11el. For a search of regions of the inRNA where collisions detroven 2 adjacent ribusones might occur, the Cansaian distribution is replaced by a unit distribution with a witth of n amino acid residues.

<u>\$</u> Average codon usage in E. Table 1

CGC 19 AOT 4 GGC 41 CGG 1 Thr ACA 4 Val GGX 22 CGG 23 AGC 23 AGA 1 CGC 18 AGA 0 ACC 23 AGA 1 CGC 23 CGC 2 AAC 20 CGC 2 AA	Jrg (15.)	_	XX.	<u> </u>	CAC	=
CGG 1 Dir ACA 4 Val GUA ACC 25 ACC ACC 25 GUC ACC 25 GUC CUA 1 ACC 2 CUC 5 GUC CUC 5 GUC CUC 5 GUC CUC 5 AAU CUC 5 AM CUC 6 GUC 14 CUC 7 AU CUC 7 AU CUC 8 AU CUC 8 AU CUC 9 AU CUC 1 CAC CUC	כמנ	<u>6</u>	305	., 88	5,45	=
CGU 33		_	Y.	čů;	C.YC	:2
MOA 1 ACC 5 GCC AGG 0 ACC 30 GCC CCC 1 Pro CCC 3 AAA CCC 2 CCC 3 AAA AAA CCC 3 CCC 3 AAA AAA UCG 4 AAA AAAA AAAA AAAA AAAA		2			CAG	١~
AGG O ACU SO CUC CUX 1 Pro CCA 7 Lya AAA CUX 5 CCC 2 AAU CUX 8 CCC 3 AAU CUX 5 CCC 3 AAU UUA 5 AN CCC 3 AAU UUG 6 GCC 14 CAG UUG 8 GCC 14 CAG UUG 3 GU 24 CAG UUG 3 GU CAG CAG UUG 3 GU CAG CAG UUG 3 GU CAG CAG UUG 4 CAG CAG CAG UUG 4		_			ပ္ပ	+
CCC 5 CCC 2 AAU CCC 5 CCC 2 AAU CCC 6 CCC 2 AAU CCC 3 AAU CCC 4 AA		0	30	GEC	:cr	*
CVC 5 CCC 2 AAU CVG A2 CCC 3 AAU CVG 3 AAU CVG 3 AAU CVG 4 CCC 14 AAU CVG 6 CCC 14 CAA CVG 6 CCC 14 CAA CVG 7 CCC 14 CCC CVG 7 CCC CVG 7 CCC 14 CCC CVG 7 CCCC CVG 7 CCCCC CVG 7 CCCCCC CVG 7 CCCCCC CVG 7 CCCCCCCC CVG 7 CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	Ġ	_	Ş	414	S	8
CUC 25 Ass AAC CUC 25 Ass AAC CUC 3 AAC CAA AAC CUC 14 CAA CUC 14 CAC CAC CUC AAC CUC 14 CAC CAC CUC AAC CUC AAC CAC		163	g	3.46	ຄຸກວ	2
CUU 5 Ale CCU 3 AAU UUA 5 Ale CCA 28 GIn CAA UUG 6 UCC 14 CAG UUG 7 UCC 14 CAG UUG 7 UCC 14 CAG UUG 8 UCC 14 CAG UUG 10 UCC 14 CAG UUG 10 UCC 14 CAG UUG 17 UCC 14 CAG UUG 17 UCC 14 CAG		Z.		3.40	the AUA' o'	•
UVA 3 Ala Octa 28 Offic CAA UVG 6 CCU 14 CAG UVG 8 CCU 14 CAG UVG 3 CAC CAC CAC		Ö	돥	4.4 U	2(1)	=
UCC 6 OCC 14 CAG C		~ 7	ź	C3.3	31.0	느
UCA 3 QCO 28 His CAC UCC 14 QCO 46 GAU QCO 3 GIN GAA UCC 11 QAO		æ	ဗ္ဗ	C.C	760	2
14 GOT 46 CAC 3 GIV GCA 3 GIV GAA 13 GOC 31 GAG	ron.	43		CAC		-
3 Gly GGA 3 Glu GAA.		z		CAC		
070 IE 200		es	3	5		
		<u>:</u>	ဗ္ဗ			

Determination of this codon usage has been carried out as described in Materials and Methods. Values are expressed per thousand and are approximated to the next integer.

latues of tRNA concentratious used in the calculation of the average number of selections (N) expressed as % TABLE 2

Amino aciel	(Lelon	-	Ë	Amino acid	Codon	2	S
Ang	(CIL), C. A)	S-125		1.3	enne (,10:+	
,	ii S	구 중) eCc	21 41	
	ઝલાંગ. બ	-29 -5			લવાય. લ	寄占	
]va	CO(C) C)	2 2		ij	1334	17	ਰ ਦ
	ra:	- 			1.3.36	<u>.</u>	1
	9 3 3 5 5	3·6·1	4.7	ne 4.	AAC. CI	3-31	
	: :: -:::	<u></u>) 6 6	Gin	CAA	₽.	
j	322	a and	:		CAG	주 구.	
ķ		7 (P)		¥	CA(C, C)	2.21	
	CON G	=		සි	GAA	50.5	3.7.5
	AGIU, C)	?			GAG	α	<u> </u>
è	l ACU.	5:01£		risk.	GAC. CI	‡	
	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	주 F		T,	เราเก	86	
ä	(AC. (A)			S.	1300 CI	L-13.	
2		, de		Phe	ממונר, כו	90·1	
	OC(A. 0)	3.68		Пв	AU(U. C)	5.63	
4	1000	3-80			,4U.A.	61	
	25	* -0-+			AUA.	0.60	
	FC(3, 6)	F-1:0	į) <u>k</u> (אונס	<u>89</u>	
Ę	ຕິດປະ. ຕາ	<u></u>		Trp	001	₹	
	r:C:A	- 					
	• • •	 Ot : 1					

Recognition pattern according to (1) themura (1981a.b); ((3f) Gresjean & Fiers (1982)

Rengaition of I cooks by 2 IRNAs was taken into occure interpolated values

Detributed salves.

Two different possible values for concentration (see Materials and Methods). Apparent concentration.

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TRANSLATION IS A NON-UNIFORM PROCESS

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mRNAs (Varenne et al., 1982), the nucleotide sequences of colicin El (Vannada et al., 1982) and colicin A (Morlan et al., 1983) have been established. Since frequent usage of codons corresponding to minor iso-IRNAs of E. coli was observed (Murlon et al., 1983), it became more likely that codon usage, rather than mRNA secondary structure, was the predominant factor in the mechanism responsible for Since our previous report of a non-uniform rate of translation for some colicin creating pauses in elongation.

polypeptide chains required the establishment of experimental conditions leading to results reflecting as accurately as possible the true concentrations of the A detailed analysis of discrete intermediates in the elongation of nascent intermediates.

(a) Detailed analysis of colicin A interprediates

·guara a pis

The main problems concerning quantitation of the asseent polypeptide chains ralies; (2) protectisis should be avoided or at least minimized; (3) the recovery of polypeptides in the solubilization process should be constant for all intermediate were as follows: (1) their radiolabeling should be as uniform as possible for all M, sizes: and (4) the yield of immunoprecipitation thould also be constant.

In analysis of colicin A intermediates, the following conditions were used.

colicin A represents more than 50% of total protein (Varenne et al., 1981) and a labelling in spite of the unequal distribution of methionine residues along the patypeptide chain. This unexpected result, which is analyzed in the Discussion. was deduced from the comparison of fluorograms (not shown) obtained from total cell proteins of the fully induced strain CA31 Cold radiobabelled with a 14C. labelled amino acid mixture or with [158] methicuine. Under these conditions, (1) Pulses of 20 to 25 seconds with [158]methionine allowed a rather uniform direct comparison of many intermediates is possible.

(Lin & Zabin, 1972). The same phenomenon could be observed for nascent chains of fluorograms of trichloroacetic acid precipitates and insmunoprecipitates of fully proportional to their length but are apparently determined by their conformation of edicin A in vitro (Varenue et al., 1981) or in viro (Fig. 1(a), lane 2). Comparison induced cells (Varenne et al., 1982) showed that with the experimental conditions described under Materials and Methods, this protectivess was generally weak for (2) It is well known that incomplete polypeptide chains made by nonsense mutant strains of E. coli are degraded at different rates, which are not directly coliciu A intermediates.

antigen ratio (an increase of this intio led to perturbations in the pattern of (3) and (4) The best conditions for solubilization and immunoppecipitation were performed (Fig. 2. lane 3), all the intermediates and the terminated colicin A were intermediate immunoprecipitation was apparent, due to a limiting antibulyintermediates and must be avoided). The low M, intermediates that excuped determined. When a very efficient solubilization treatment of soil membranes was present in the immunoprecipitation mixture, but considerable inhibition of low M,

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colubilised as described (Navenne et al., 1992) and a second flanz 4) was solubilized as indicated in blaterials and Methods. Mahemlar uvigith standards: bovins serum albumin, US,000 M; orallomin, 46,000 M; carbonic anhydrate, 20,100 M; serbeat trylish inhibitor. 20,100 M; bysogyme, 14,400 M. protectivitis and solubilitation terbaique in recovery of collein A intermediates vas added. One sample (kane i) was immediately solubilized and immunoprecipitated; another (fane was first incubated at 37°C for 1 h, then sulmitted to the same treatment. One sample fully induced cells were pube-tabelled for 200 with [1935]methionine and chloramph

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efore inmanopreceptation either (fine 4) as described (Varenne et al., 1982) or (iam. 3) as indicated in Materials and Methods. The ougernatent of the fatter incountegrecipiane was entimited to an additional amountsprecipitation (large 1). Peket recovered ofter the milk solubilization was evolubilized mendence in the colubalization process. Pully influed cells were pulse-inhelike for 200 and solutifized through the harst peruse and impartopirecipitation was upuly carract out flane II. 2. Consequence of hinding of large intermediate mesent polypeptible

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immunoprecipitation could be recovered in the supernatant and were totally immunoprecipitated in a second step procedure (Fig. 2, lane 1). Conditions that cause an incomplete solubilization of the membrane fraction (Fig. 2, lane 4) allowed an efficient immunoprecipitation of low M, intermediates because much less competition by mature colicin A then occurred. Missing intermediates of high M, and colicin A could be recovered (lane 2) from the membrane pellet with a harsher solubilization process similar to that used previously (lane 3).

harsher solubilization process alonilar to that used previously take of.

By combining densitemeter profiles from fluorograms obtained under both conditions in another experiment (Fig. 1(b), lanes 3 and 4), corrected for low M, intermediates by quantification of these intermediates in whole fully induced cells (not shown), a profile was obtained that reflected the real intensities of intermediates (Fig. 5(a)).

Thus, experimentally we have access to t, the mean duration of addition of a given acid residue corresponding to a given codon. If the tRNA concentration in colicin A-producing cells were known, it would then be possible to check if the non-variable rate of elongation might be explained by the stochastic model described in Materials and Methods.

(b) The internal pool of tRNAs is not perturbed upon synthesis of colicin A

Intracellular tRNA concentrations for E, cofi have been determined by various authors and particularly by Ikemura (1981a). In order to use these tRNA concentrations in our calculations, we had to make sure that the internal pool of various tRNAs was not perturbed upon synthesis of a very highly expressed protein like colicin A that displays an unusual tRNA usage as shown in Figure 3. To allow an easy comparison with other proteins examined by Ikemura (1981a,b) in E, coli, the data points were analyzed by linear regression. The regression line is expressed by y = ax + b. The amount of tRNA = x, the frequency of tRNA usage = y, the correlation coefficient = r:

with 19 1RNA concentrations, $y=4\cdot2x+0.80$, and r=0.65, with 23 1RNA conventrations, $y=3\cdot7x+1\cdot12$, and r=0.63.

These results compared with those of Ikemura (1981 α) clearly show that tRNA usage of colicin A is different from highly or weakly expressed proteins, and more generally different from the mean usage of E, roli proteins (see Fig. 3), for which y=6.9x-0.48 and r=0.96.

As previously emphasized (Varenne et al., 1982), an increase of about 100-fold in colicin. A synthesis by induction had no effect on the intermediate intensities, but mitomycin C only increased the number of induced cells and did not modify the amount of colicin. A produced by each induced cell. Therefore, this stability of intermediates after induction did not allow any conclusions to be drawn as to the effect of the amount of colicin. A produced in each cell. In order to clarify this point, rells were pulsed and chased at different times after mitomycin C addition (Fig. 4, hance 1 to 9).

Under the conditions used, all cells were induced after 15 minutes of incubation

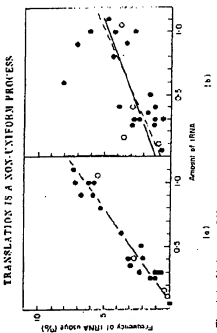


Fig. 3. The relationship between tRNA abundance and its usage found for E. coli genes and for one (gene of colicins A). For the reasons indicated by Nemara (1981a), data for tRNA, A, Abl. The 4-1 and Set are not shown. Data for Girl. Girl. St. Vall and Vall are specified by open circles (see the text). Linear repression analysis was performed with 23 tRNAs (continuous linea) and with only 19 tkNAs (or colicina A (trosten line) to deciliate comparison with individual proteins analysed by tkNAs (1881a).

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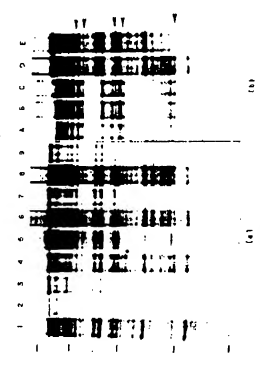


Fig. 4. Intermediates are not incified by overproduction of colicin. A. Positioning of intermediates.

(a) Cells were pulse-labelled either before induction (lance 2 and 3) or after inclinities of 15 min (lance 4 and 5), 55 min (lance 4) and 5), we lost min flaires 8 and 9). In lance 1, the sample applied was similar to that applied in lance 1 and 5), were interminantly independent allow detection of the mage of chare. Dis Cells were inte-labelled at 35°C with (Philippinoine for 4 a flaire A), is than 19; 8 without 1. Induction of the mage of chare. Dis Cells were interiorably the chare of 30 s and 603, respectively. Arrows indicate intermediates just downstream from wellinmine residues 183, 202, 195, 387, and 448.

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with initioniven C, and the intermediates observed (lanes 4 and 3) were essentially produced by newly induced cells (compare to lanes 2 and 3, which show coform production by induced cells). After 35 minutes (our routine induction time; lanes 6 and 7), tenfuld more colein per cell was produced than after 15 minutes of induction and about fivefold more was junctured after 100 minutes of induction and about fivefold more was junctured after 100 minutes of induction and about fivefold more was junctured after 100 minutes (lanes 3 and 9). In all cases, arcumulated interactifiates were located at the same If, and had the same relative concentrations. The artefactual differences observed from top to bottom between lanes 4 and 6 may be explained by the low yield of immunoprecipitation of low II, intermediates when amounts of complete colicin A become too large, as shown in Figure 2. Two lines of evidence demonstrate this point: (1) the chase was similar in lanes 5, 7 and 9; (2) when a milder cell solubilization method was applied to cells induced for 55 minutes, profiles obtained after 15 minutes (lane 4) and 55 minutes (lane 1) were directly recovery of intermediates in the solubilization process was equivalent.

This experiment clearly indicates that accumulations of intermediates (reflecting "pauses") routinely observed when colicin A is highly expressed do not result from perturbations in tRNA concentrations induced by the overproduction of colicin A. Furthermore, the experimental results in which the Citrobaster freundii strain CA31 ColA was need could be compared with theoretical predictions obtained based on E. coli tRNA concentrations, since we observed the same pattern of intermediates when the plasmid pColA was introduced into E. coli K12 W3110 (Varenne et al., 1982).

(c) Correlation between theoretical and experimental prafiles for colicin. A

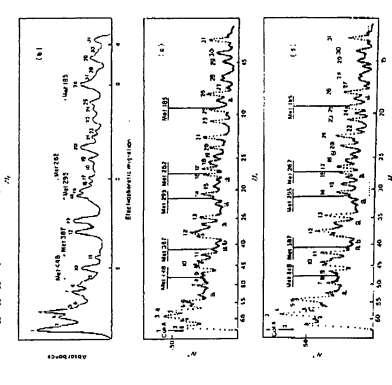
The theoretical profile corresponding to t=AN+B cersus the position of intermediates could not be plotted, since A and B are not known; however, it is possible to plot N. Then, maxima and minima of N correspond to maxima and minima of I and, if the model was correct, must correspond to maxima and minima of the experimental profile. Indeed, the amount of maseunt polypeptide chains comparising a amino used residues at the P-site of the ribosome is proportional to the mean chustion of addition of the n+1 amino acid residue.

During to dispersion (see Materials and Methods), correlation was sought between the scans of intermediates (Fig. 5(b)) and the plot of N* (the dispersed values of N: see Materials and Methods) as a function of electrophoretic migration. The most visible peaks of the fluorogram were numbered from the function A) to 31 (last visible intermediate).

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We observed that it was not possible to optimize the dispersion of N (a part of the N plot is shown in Fig. 6) over the whole range of polypeptide sizes, even by using a constant dispersion in M (ΔM_i = constant), and two different dispersions N^* with different full width at half height with Δx = constant had to be used. The best it between theoretical and experimental profiles was obtained with the dispersion shown in Figure 5(c) for the high M_i range, and in Figure 5(a) for the high M_i range.

Since use of the experimental value for tRNAII concentration resulted in an



(1982). The autobers of the amino acids (Markat of etc. 1983) everysombling to the topo of the unin beoretical peaks in the N* profile are indicated in parenthese, 2 1537, 3 (519), 4 (533), 5 (518), has been protted as a function of pulypeptide chain. M, with a field width at half maximum (FWHM) = 116 mm, The distance between peaks 1 and 31 was 5811 mm, (b) Densitonates profile for rolicis). I intermediaten, Cella were pular-lahellad for 301 aund entarged Hammyranus from त (अपने , र (मेंदर), ४ (मंदर), १९ (अपने), १९ (आपने), १५ (अपने), १५ (अपने), १५ (योग), १५ (योग), १५ (योग), १९ १७ (योग), १५ (योग), १९ (योग), १९ (योग), १९ (योग), १९ (शोग), १९ (एस), १५ (एस), १५ (एस), ig. If it were scanned free the text for details). Electrophysetic migration was foun right to left methionines in positions 14%, 38%, etc., are located between the inclinated junition and the next Fig. 5. Conquirison of theoretical and experimental profits for intermediates. (a) Nº (see Materior Letters a. b. e and d indicate migrations of calibration process (a = 14.1811. 1), b = 20.1881. U, indicate the upstream interrrediate (not vinible in the pulse but visitle in the strass), (c) Same as (a) EWHM = 6-4 mm (d) Same as (c) but the revoluing spectrum of (ENS) was that of Cosjona & I Arrowhends: P. Met 148. P. Met 185. cle. and 4 = 48,000 14). 38 (360), 20 (1 Hd), 30 (1 Hr), 31 (131) and Methodel J. 00000 = 3

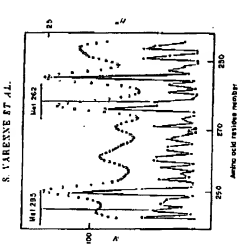


Fig. 6. Plotting of X and X* for part of the colocia A sequence centred around smine acid residue 170. In the N profile: (\$1) computed from experimental RRNA concentrations; (O) computed from interpolated RNA concentrations. The N* profile is a part of that shown in Fig. 3(c).

N value of 345, and for N^* profiles caused accentuated peaks at positions 255, 426, 533 and 567 (not shown) that did not have an equivalent counterpart in densitometer profiles, we suspected that the experimental value could largely be underestimated. In contrast, when the interpolated value was used (then N=167), the peaks cited above were more equivalent to experimental peaks. This value was therefore used routinely for further calculations.

Although the observed theoretical profiles A and C do not exactly reproduce the experimental profile B, there is generally a good correlation between positions of theoretical and experimental peaks. It is also quite significant that regions where N** is low [between peaks 13 and 12 and between peaks 13 and 14] contain a lower amount of intermediates than ekewhere. Some minor theoretical peaks (0a, 11a, 11b, 13a) correspond to slight bends on the fluorogram, one (25a) does not appear in this experiment but is just visible in some others, two (12a and 15a) are never observed. In contrast, the experimental peak I does not appear in the theoretical profiles, since it corresponds to the colicin A itself and not to an intermediate, and the experimental peaks 22 and 24 are not predicted with Hemura's (1881a) recognition pattern. As this pattern was slightly different from that reported by Grosjean & Fiers (1882), see Naterials and Methods, it was important to check the effect of this difference in the above correlation. In the new pattern obtained (Fig. 5(d)), the position for most of the peaks remains unmodified although areas are locally changed. The main difference can be observed for peaks 22 and 24 that now somes.

This feature is particularly interesting, because the modifications in these two leaks result only from the existence of three GAALIAGI-GAA codous for glutamic acid in positions 206-207-208 for peak 22, and one GAG codon in position 190 for

peak 24. The better correlation for peaks 22 and 24 suggests that indeed two different tRNAs may exist for glutamic acid instead of one. The effect of this change in decoding pattern for GAA and GAG was checked elevener in the profile but no clear supplementary evidence favouring this proposal could be put forward, because other modifications do not appear in regions where N° is low. For the same reason, no conclusion could be drawn for the four other codons. Repeated similar analyses for other proteins, with the eventual help of site-directed mutagenesis and insertion of oligonucleotides, could be useful for the clarification of these ambiguities in decoding pattern.

the should be pointed out that the molecular weights of some peaks in the densitometer profile A were accurately determined in the following way. Very short, [135]methionine pulses (Fig. 4, Ianes A, B and C) were performed in fully induced cultures of CA31 ColA strain. In regions of the gel where methionine residues were close enough, all intermediates appeared. But, in other regions, only the intermediates immediately downstream from a methionine residue could be seen, while intermediates just upstream could not. Thus, it was possible to determine accurately the real molecular weights of intermediates near methionine residues 185, 262, 295, 387 and 448.

From this comparison between profile B and the profiles A, C and D, the following conclusion can be drawn: in spite of technical difficulties and theoretical problems exposed in the Discussion, the marked correlation that exists between positions of peaks in theoretical and experimental profiles indicates that accumulations of mascent polypeptides are indeed directly related to the tRNA concentrations. This conclusion will be discussed further after analysis of other non-uniform translations.

(d) Intermediates in synthesis of colicin EI

The endon usages for the colicin BI gene (Yamada et al., 1982) and the colicin A gene are rather similar (Morlon et al., 1983); thus similar experimental results were expected for translation of colicin BI mRNA. Marked intermediates were in fact observed in a pulse-chase experiment (Lazdunski et al., 1984), and the correlation between position of the Rheovetical and experimental peaks was checked. However, owing to the small number of methionine residues in the protein and to a poor yield of immunoprecipitation in the low M, range, this correlation is more difficult to establish firmly, and a numbered correspondence between peaks cannot be proposed.

(e) Intermediates in synthesis of TEM 1-8-lactomose

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The phenomenon of non-uniform translation was investigated in race for some ather proteins in our previous work (Varenne et al., 1932). The theoretical approach described for colicin A was applied to these proteins: TEM 1- \$\beta\$-lactamase encoded by pBR322, the OmpA and LamB proteins, and the ekugation factor EF-Tu. Theoretical profiles indicated that marked intermediates could be expected to orcur in TEM I-\$\beta\$-lactamase. As intermediates were not

experimental and theoretical peaks confirms that pauses are not created by an imbalance in the puol of the tRNAs, but are related to the physiological especially in the low M, range, the correlation observed between positions of Although the intermediates are more difficult to observe than for colicin A, differences in conventrations of the tRNAs.

It must be pointed out that if the experimental concentration reported by for which only the interpolated concentration of this IRNA could be used. This last result suggests that the Cold plasmid might modify the intracellular concentration of this tRNA. If true, this would occur in a permanent way, since kemura (1981a) for the tRNAmes, was used in computations, a reasonable fit described in Figure 4. Further experiments are nearly to check this passibility no modification of the pattern of pauses could be observed in the experiment could also be observed (Fig. 7(c)) in contrast to results observed with colicin A, and to evaluate more accurately the concentration of tRNAmics in E. coli strains.

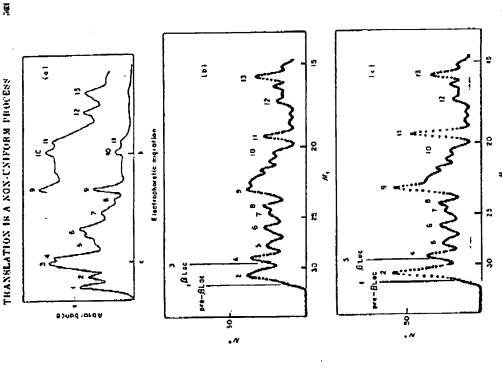
(f) Internediates in synthesis of pre-Ompal protein

7 × 10° copies/cell) and OmpA protein (about 10° copies/cell), use of minor tRNAs is avoided. Nevertheless, the other tRNAs do not exist in equal amounts: the ratio of the two extreme IRNA concentrations used in synthesis of these proteins is (Varenne et al., 1982), the synthesis of this last protein was analysed and the two In exuclesies of constitutively highly expressed proteins like EF-Tu (about concentration taken as the interpolated value, and 22 with the value from protein in Figure 9(b). Since a possible secondary atructure for the mRNA of Ikemura (1981a). Faint intermediates would be expected, as shown for the OmpA OmpA has been proposed (Movva et al., 1980), and since the possibility that hairpine might be involved in discontinuous translation of OmpA was suggested adenut 5 instead of about 10 for TEM I-G-lactamase with the IRNAMIN hypotheses concerning discontinuous translation were examined.

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Discontinuous translation was again observed with some differences in profile as compared to A typical pulse-chase experiment is shown in Figure 8. those for rolicin A and TEM I B-lactamase.

translation was uniform in this region; in fact, for an unknown reason, a major (!) Pauses were never observed abore 23,000 M, but this does not mean that hart of the prowing polypeptide chains was lost in all similar experiments, since a



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according to livenara, 1981a b.1 (a) Ibrasionarter profile d interraciates. The upper trace corresponds to the saming of the same gel but exposed 8-fold longer. (b) 3° with FWIM) = 11 d inm. (c) Same as (b) but the experimental correctation of tRNA_{lliver} was used. § Lac and pre-files. § stace and pre-glactanists. Fig. 7. Comparison of theoretical and experimental profiles for eta-bectanase. (Recognition spectrum

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observed. In any case, interpretation of the upper part of the profile would be mission translation should produce a continuous background, which was never difficult for two reasons: first, the abnormal migration of mature OmpA (Nakamura & Mizushima, 1970) and thus propably of high molecular weight intermediates could not be avoided totally; and second, a part of the precursor

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form is containable formally processed when the size of the nascent polypeptide chain becomes greater than 30.000 M, (Josefsson & Randall, 1981). Thus, four differently migrating forms are expected for each kind of intermediate.

(2) As expected if pauses are due to IRNA availability, observed accumulations in OmpA synthesis were less marked than in codicius or TEM 1. Flactamase synthesis, and a much longer exposure of the fluorogram was needed to detect the pauses shown in Figure 10.

(3) Consequently, the background was more important (Fig. 8, Inne 2), and must be taken into account for interpretation of the densitometer profile of lane I shown in Figure 9(a) and (c). Some bands that do not disappear during the chase, and are also visible in the same chase experiment followed by immuno-precipitation with anti-lipoprotein (lane 4), correspond to abundant proteins of the cell (lane 3) and must be discarded for the analysis of the experimental profile of Figure 8(a) and (c).

Theoretical profiles of N* are shown in Figure 9(b) and (d). Amino acid residues whose numbers are indicated on the abscissa of Figure 9(d) are those which are bound to tRNA in the ribosome A-site just upstream of the ribosome entry into possible hairpins of the mRNA (Movvu et al., 1980). A strong correlation again exists between positions of observed and predicted peaks in the tRNA theory, whereas a full corvespondence does not exist in the hairpin theory. Moreover: at least there peaks (2, 3 and 6), curresponding to residues 206, 192 and 164, are found in regions of mRNA where non-optimal codons or non-classified codons (corresponding to His. Asp. Cys or Ser: Ikemura, 1981b) are not encountered.

Two conclusions may be drawn from the above data. First, even for constitutively highly expressed proteins, faint intermediales (which reflect the unequal duration of searches for adapted ternary complexes for the cockon in the

Fig. 8. Intermediates in pre-Omph synthesia. Cella were pulsed for 30s with (**5)methinume (fame) I and chased for 12ths (fames 2. 4 and 5), then notabilitied and framunoprecipitated by anti-comph protein antibody (lance I and 2) or by anti-fipoprotein (lance 3 and 4). Lanc 5, whole cells.

<u></u> 3 = 9 TRANSLE TO NOT A NOT LEAD TO SEE THE Etectropharent migralian Electrophoretic mgration 2 ₹ 25 3 ន 202 #2000010£4 6 40u0q/01Q4 33 . N . N

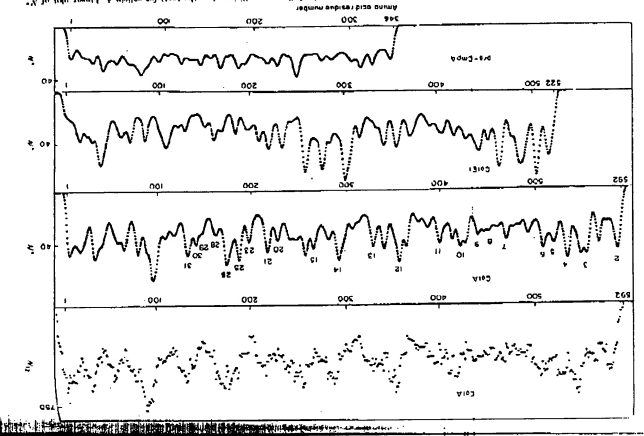
Fig. 9. Camparison of theoretical and experimental profits for pre-funged. (Recapitation spectrum according to Remara. 1981a.b.) (a) Bensimmeter profits of intermediates obtained from has 1 (Fig. 8). The filled circles indicate the bands that do not disappear during the chase exist is those 3.4 and 5 and do not constitute intermediates of synthesis. (b) N° with FWHM = 8.4 mm. (c) Partial enlargement of (s), (d) Tartial enlargement of (b) The numbers indicated on the abactian correspond to the amino acids greeified by the codon exposed in the ribbonous A site just upstream of the ribconne entry into passible latigates in the mRNA (see the text). The numbers of the mainto and sceneral profits to the tops of the theoretical packs in the N° partile are indicated in parentheses. I (218), 2 (2001), 3 (102), 4 (184), 3 (172), 6 (104), 7 (154), 8 (147).

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Asite of the ribusomel can be observed even in parts of inRNA where only "optimal" codons are used. Secund, the possible role of inRNA sexundary attructure in creating a non-uniform rate of translation is probably a minor one, if it exists 81 all, for the OmpA protein.

Fig. 19. Configuration between theoretical profiles, From top to letterm; search of these collisions (see the text) for colloin A, Tarent plat of X* with FWHM = 5.8 unine acid residues for colloin A. Linear plat of X* with FWHM = 5.8 unine acid residues for colloin A. Linear plat of X* with FWHM = 5.8 unine acid residues for pre-OmpA profiter. Recognition spectrum according to Ibernal of the comparation.



(or N.) presents certain particedurities along the polypeptide chain, Observation of the theoretical profiles presented in Figures 5. 7 and 8 does not ensily provide this information for two reasons: (1) the X-terninal parts of the differences in aminu acid residues. If, values and by logarithmic migration of It was, therefore, more appropriate to plot $N_i = f(x_i)$, where k_i is a linear proteins are lacking; (2) distartions in the profiles of N* are introduced

polypeptides in the gel.

Since the rate of translation is lightly sonnected to N. it is of interest to know if

(B) Computeson of No turnations along different mRNAs

function of the amino acid residue number and to disperse the results. Three Y^ullet important extent, local variations (see Fig. 6). As mentioned above, the ratio of profiles are shown in Figure 10 in order to compare two proteins sharing similar though very efficiently expressed (OmpA protein). It must he kept in mind that view of the slowing down regions along the whole mRNA, but minimizes, to an the two extreme values of V is 10 for both colicins. and 5 for OmPA: 3% of OmpA features (colicins A and Ell, and to another that is very different from both, the choice of the N. representation instend of that employing N allows a general codons and IT.'s of colicin A codons lead to Whigher than four times the minimal ralue of N (N minimum = 16.3).

(1) Regions with a high number of tRNA-codon interactions exist along the whose polypeptide chain and not mostly in the C-terminal part of the molecule, as might be suggested from previous theoretical profiles plotted in a semi-logarithmic representation. Pauses under 13.500. M, were not detected in whole cells of fully intermediate polypeptide chains in this range of $M_{
m t}$ were probably very sensitive indured CA31 CalA. Two reasons at least could explain this observation: first, to proteolytic degradation: secondly, distances between two adjoining polypeptides are approximately threefold smaller below 14,000 M, than above (Swank & Munkres, 1971). Since the dispersion in the gel remains similar, Several remarks can be made about the .V. plots shown in Figure 10. interference is much more marked between close intermediates.

(2) The variability of N* during translation of mRNAs is much higher for (3) No, the average value of No calculated over the whole polypeptate class. is colicins A and El than for the OmpA protein.

higher for colining A and El than for the Uniph protein. This reflects the

Sum of N for the whole protein Number of amino acid residues

As a close relationship between high expressivity and low values of N was found by Gouy & Cautier (1982), values of S were calculated with two decoding patterns (from Ikemura. 1931a, and from Grosjean & Fiers. 1982) for the proteins studied in this work (Table 3). To facilitate the discussion, this Tuble also contains further information; (i) differences in F from protein to protein may result, at kast partially. Irom differences in araino acid composition. To take this

among these proteins.

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difference between N values:

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Tulnes of the average number (S) of selections for some mRNAs TABLE 3

1									Average.
Leroding pattern	3	Cale	Ē	Отра	Leng Bar	PhoE	Tuda	LANZ	S. cali
					١	1	-	0.0	1:50
_	3,6	1.23	7 2	30.8	?	2	2		7 9
ن.	700	7.80	X.	25.3		Ţ	2	in N	0.7
5 5	9	9	1	6	ž	į	61 6	Ş	60 61 61
ž -	•	•	:	}	. :		•	4	<u>!</u>
<u>وَح</u>	ㅈ	ß	æ	æ	2	Ŗ	~	3	=

In calculation of .S. the interpolated value of the tRNA_{line} concentration was used. Deriding pattern. I. according to Ikemura [1981a,b]; OF, Greatern & Fiers (1982); Io, Ikemura optimal codon

(ii) PloE, and figalactosidase are included; (iii) the everage tRNA usage from Table ! allows estimation of N for the total B. coli cell proteins. These estimates using for each amino acid the codon(s) corresponding to the major iso-tRNA; point into account, the minimal N value was calculated with Ikemura's decoding nattern. This optimal is value corresponds to an optimal nucleotide sequence are also included.

[1978], respectively. In our program, different values of a between 11 and 17 were used in the theoretical treatment for colicin A. For each codon i, the computer maximum value, especially if such a value has not been attained upstream in the mRNA. Thus a diffusion computer program, where the Gaussian distribution was motein synthesis presented by Bergmann & Lodish (1978) or Van Heijne et al. calculates the total average number of trials relative to the next n codons just In being the minimal possible distance between two adjacent ribosomes) reaches a replaced by a unit distribution on a codons, was applied to N versus residue number. Minimal distances of 12 or 15 codons were used in the kinetic models of mRNA where the total average number of selections relative to madjacent codons ujestream rabosome may also pause because its movement might be impeded by the first. In the model for translation analysed here, regions where such a phenomenon would have a maximum probability of occurring are the I coding extremity of the mRNA if the termination rate is limiting, and in parts of the (4) If ■ ribnsome pauses too long in a specific region of the mRN.4, the next downstream from codon i (i.e. from codon i+1 to codon i+n).

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"secondary pauses" were not detected in scanning intermediates, even in regions corresponding to maximu of the N_{12} plot. It is of interest that the 12 first endoms width at half maximum = 5-8 residues), and particularly that the most important one was located, as in the N^ullet profile, around residue number 95. The other values lead to a high value of N13, which may interfere with the initiation rate of We observed that maxima of such a plot dispersed with n = 12 (designated as N_{12} in Fig. 10) could be predicted from the \dot{N}^* profile shown in Figure 10 (full of u led to similar conclusions about positions of the maxima. In fact, such translation for colicin A mRNA.

4. Discussion

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A-site of the ribosome. The next two steps in the cycle, that is transpeptidation and translocation, account for a much shorter time than the delay before the successful collision with the specific iso-tRNA; (2) insanuch as tRNAs are not in equimolar concentrations in the cell cytoplasm, including those (the most abundant) that are preferentially used for highly expressed proteins of E. coli, the elongation of polypeptide chains must occur at a variable rate for all E. coli Two important conclusions can be derived from our results (1) the rate-limiting step in the elongation eycle of polypeptide chains is the search for the ternary complex (animoncy |-tRNA bound to EF-Iu and CTP) specific to the codon at the

polypeptide chain. The second dealt with the possible loss of material before methionine residues, which might result in heterogeneous labelling along the precipitation that might not be constant for all intermediates. The most serious slways he suppressed. This difficulty appeared especially in TEM 1.8 lactamase luture use of protease mutants such lon (Grossman et al., 1983) and for the use of a We had to overcome a number of technical difficulties in order to interpret our experimental data. The first was related to the beterugeneous distribution of immunoprecipitation. The third problem came from a vield of immunodifficulty was a possible protecytic degradation of intermediates that could not experiments, and seemed to be highly variable from protein to protein. By shortening each step when possible, we could alleviate this problem. However, protense inhibitor might further alleviate this problem.

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All the difficulties evoked above were circumvented for colicin A. However, theoretical problems remain.

(1) The logarithmic migration for intermediates can suffer considerable local deviations (see colinin A intermediates. Fig. 5).

(3) Experimental (RNA concentrations are known with a non-neglectable (2) A constant dispersion was used for computation, although the dispersion along the gels is clearly not constant.

(4) The decoding spectrum is not known with certainty for a certain number of codons, and the determination of "apparent concentrations" when a codon is partially recognized in 11170 by a second IRNA is questionable because of fact of standard deviation, especially for minor tRNAs.

(5) The accuracy in concentration for those cases obtained by interpolation might be lower than that for those which have been determined experimentally for two reasons: (i) codon usage, although it was established from 62 genes, can only be approximate; and (ii) an exactly linear relation is not likely to exist between frequency of tRNA usage and amount of tRNA. The following example gives an idea of the uncertainties introduced by interpolation. According to Remura (1981a). IRNATE and IRNAM have the same concentration, but Table I indicates that their usage is quite different. Thus, interpolated values of concentrations (1-13%, and 2-33%) would lead to numbers of discriminations 188 and 43) very different from that (60) deduced from the experimental adequate data obtained in viva. **有不能的地位的一种,他们也可能是一个**

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TRANSLATION IN A NON-UNIFORM PROCESS

(b) Differences between tRNA frequencies and operational frequencies of

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(7) N_i was calculated as M_i but the probability of transpeptidation after each collision between a rodon and a cognate-tRNA is probably not 1 for each tRNA, and might differ from one tRNA to another; this would introduce increases of M_i and distortions in the profile if the perconage increase is not the same for all codons.

(8) The mean duration for the addition of an amino acid corresponding to a given codon may be obtained from N by the simple relation $t = (\theta_0 + \theta_1)N + t_2 + t_3$ only if θ_0 , t_3 and t_3 are the same for all vodon species $(\theta_1$ is independent of the codon). In fact, this is probably approximate. For example, any attempt to explain codon usage should involve both IRNA concentrations and a consideration of the energetics of codon-anticodon pairing; one tRNA can often translate two codons and a bias in the codon usage has been observed (Grosjean & Fiers. 1982: Grantlam et al., 1981: Ikemura, 1981a.b). This bias might play a role in fidelity and/or in the rate of translation of codons. There might thus exist an effect of codon choice on translation rate by modification of θ_0 , t_2 and t_3 that we could not take into account for lack of information.

Experimentally, the role of codon choice could not be demonstrated in colicin A in a quantitative way. For example, non-optimal codons (which, as a general rule, are rarely used in constitutively, highly expressed genes) with a low energy of interaction are found between peaks 13 and 14 in positions 311 and 314 for AAT (Asn) and 313 for ATT (He). These evdons do not seem to induce an additional slowing down of ribosomes in this region (see Fig. 3), but it is not possible to draw general conclusions about these codons from one example. Elsewhere, we cannot rule out the hypothesis that non-optimal codon usage leading to high energies of interaction might have resulted in additional slowing down of ribosomes.

These theoretical and experimental difficulties probably explain why areas of experimental and theoretical profiles do not match exactly, even for calcin A.

Tables 4 Applies and various professes in E. coli

Pretein	કર	Rate A (araino acidose)	Procein	بنا	Rate B
EP. Tu Ompel procein Ompal procein Lamb procein mmpe Plactonease Tem 1-2 lactonease		15-3-20-4 12-1-15-2 14-6-17-8 <9-5 7-4-9-6	EF-Ts Sy (ribosomal) EF-Ts EF-O dor repressor Test 1.8 decrance	4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	ம் அக்க்க் ம் அக்க்க்க்

The rates of assembly of mains acids were assayed; rate A, at 37% by descison (1982); rate B, at 24.5% by Preferent 1983). A was computed as in the first line of Table 3. References for makenticle expresses of proteins presented above are eised in the text. The value for PF-6 is approximate, since the makenticle expresses is incomplete.

However, a very significant overall correspondence was observed hetween positions of peaks, which indicates that experimentally observed variations in cloragation rates have their main origin in tRNA availabilities. Moreover, the importance of the gap between maxima and minima in the experimental profiles means that the value of B in the equation t = AN + B is low compared to AN. This provides a direct demonstration that transpeptidation and translocation steps occupy a short or neglectable time as compared to the aminoacyt-RNA selection step. This conclusion is strongly supported by the experimental determination of rates of translation for different mRNAs performed by Josefsson (1982) and Pedersen (1983). The marked decrease of the rates of translation observed when N is high $\{ampC, \beta$ -lactamase, TEM I- β -lactamase, as translation eyele.

However, we cannot exclude the possibility of a modulation of a rate of translation by the energetics of codon-anticodon pairing. Furthermore, additional factors like mRNA accordary structure might also contribute to discontinuous translation for proteins such as MS2 coal protein (Min Jou et al., 1872; Chaney & Morris, 1978) that we have not studied.

Experiments on discontinuous translation may lead to valuable new or supplementary information concerning translation in riso, concerning, for example: the recognition pattern of certain 1RNAs: tRNA concentrations; the possible effect of codon-anticodon energies of interaction; the possible influence of codon context. Site-directed mutagenesis and insertion of oligonocleotides should

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allow a more direct approach to these problems.

It is necessary to enlarge the conclusion that the elongation of polypeptide chains occurs at variable rate for E. cot proteins. This means that for one mRNA species and for one given codon, the duration of addition for the corresponding amino acid residue fluctuates around an average value. From this codon to the next one upstream, this average value varies in a ratio rather similar to the inverse ratio of the IRNA concentrations curresponding to these endons. Thus, the average rate of assembly of amino acids for a given protein is approximately proportional to the inverse of the average value. In old lie selection numbers for the whole protein, but individual rates for walividual mRNA vary (i.e. each individual mRNA is not translated at the same rate for a given protein. This implies that comparison between rates of translation for two proteins must be performed for the same parameter: average rate, minimal detectable rate, maximum detectable rate.

At least four experimental observations argue for dispersion of translation rates in individual mRNAs.

(1) In the experiment shown in Figure 4 for determination of the exact location of some intermediates, radiolahelling of intermediates after a 30-second clase is shown in lane D. If all individual translational rates were identical, the nascent chains that were upstream from methionine residue 202, for example, at the legioning of the chase and were not labelled (appearing as a blank area in lane C) should be longer 30 servings later, and mignate in the gel like publiciples of 300 for 400 residues. In fact, radiolabelling was present in this area, indicating

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differences in individual rates of aming acid assembly (as judged by radiolabelling attensities, delay in chase could not alone account for the observed result).

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labelling of intermediates in synthesis of colicin A rather similar to that obtained with a mixture of 14C-labelled amino acids, in spite of an irregular distribution of methionine residues along the polypeptide chain. The duration of pulse labelling could not alone account for this fact if the clongation rate was the same for all (2) As mentioned above, a [35S)methionine pulse for 20 seconds provided

(3) In colicin El experimenta, the translation rate deduced from the since the C-terminal methioning residue is in position 370) is higher than the translation rate deduced from the disappearance of pauses (not shown). The same Appearance of [138] methionine in mature colicie El (calculated on 152 residues, conclusion can be drawn from the colicin A experiments.

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(4) For B-galactosidase (N = 38 as for colicin Al, a variable rate of translation,

This dispersion of individual rates of translation explains why the minimum ranging from 8 to 15 amino acids per second has been reported (Talkad et al.,

induction. It is significant that the minimum values for colicin A and g-galactusidase are similar, as both proteins have the same N value, and similar ralues for the P, index from Gouy & Gautier (1982), which characterizes the 1982) must be compared with the minimal value for $oldsymbol{eta}$ -galactosidase, and not with the maximal value deduced from appearance of enzymatic activity after choice between codm-anticodon pairing energies (048 for colicin A, 043 for detectable translation rate deduced for colicin A (592 ammo acids translated in about 70 to 80 s leading to an approximate value of 8 residues(s: Varenne et at., 8-galactosidase).

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In agreement with previous studies (Gony & Cauties, 1982), the two constitutively highly expressed proteins in Table 3 have an N value close to the optimal value, since deviations between N and N optimal are 3% for EF. Tu and 6% for OmpA protein, in contrast with other proteins in this Table (35% for P. galactosidase, 51% for culicins A and El, 64% for TEM 1. B. lactamase). This genes highly adapted to the tRNA content of the cell for fast translation. It is of interest to observe that the degeneracy of the genetic code introduces an tRNAs corresponding to each amino acid were exclusively used in synthesis of attain values of 63 and 03, respectively; i.e. about 2.7-fold the minimal possible value (about 3.3-fold the minimal possible value if the recognition pettern are used in the calculations). Among known nucleotide sequences of proteins confirms once more that constitutively highly expressed proteins are encoded by solicin A or OmpA protein, for example, the average number of selections should according to Grospan & Fiers (1982), and the experimental value of tRNAmics sinthesized in E. coli, the highest values are never approached, in contrast to the lowest values. The highest value among known nucleotide sequences is 61.2 for important potential variability in possible N values. If the less abundant isothe intraunity protein for CoIEI (maximum possible ralue, 77:3: Lloubes et al.,

One can address the question of the physiological significance of discentionous:

and that the essential objective of more or less marked general slowing down is to concentrations in the cytoplasm or whether it reflects any particular regulatory strategy of the cell. It is now well-established that highly expressed mRNAs generally use abundant tRNAs and "optimal" paining energies, and that neakly expressed mRNAs often display an opposite choice (Grantham et al., 1981; Grosjean & Fiers, 1982). leading to a slower translation. It is quite passible that lucal variations in elongation rate along mRNAs have no physiological finality. introduce a constitutive modulation in average rates of translation for the mRNAs, aerording to cellular needs (Gony & Gautier, 1982). Besides, since transcription and translation are coupled, it is possible that transcription also occurs at different average rates for different genes and that the use of rare isotRNAs merely edapts translation to slow transcription. The fact that the same intermediates were also observed when transcription was blocked (Varenne et al., 1982) does not exchade the possibility of the existence of such an adjustment between rates of transcription and translation.

translation rate might be part of a tight coupling mechanism between However, local variation in translation rates, at least in specific cases, might have a physiological significance, for example by favouring sequential polypeptide medium-range interactions to take place before long-range interactions in the polypeptide chains, thus favouring domain formation. With regard to this point, 1978; Olino-Iwashita & Imahori, 1980). Furthermore, tight coupling between transcription and translation might be necessary or advantageous to ensure efficient synthesis. Discontinuous transcription, which has been reported in Chamberlin, 1981), might be a more general phenomenon, and variations in chain folding. Variable rates of polypeptide elongation might allow short and it should be recalled that colicins have well-structured domains (De Graef et al., specific cases (Darlix & Fromageot, 1972; Vanofsky. 1981; Kingston & transcription and translation.

These problems bear on molecular hiology but also obviously bear on biotechnology. It is desirable to use a microorganism having a tRNA pool as much adapted as possible to the message being translated, or if the gene is an significance of the phenomenon described is to globally attune syntheses; this artificial one, to use only optimal codons from the host organism, if the only should allow highly expressed proteins to be produced as rapidly as possible, andfor as faithfully as possible andfor as economically as possible. However, efficiency of prochection of a profein is not always related to an uptimal codon usage. With regard to this point, the case of β -galactosidase is particularly similar situation exists for colicin A, for which a very high level of synthesis lunpuhlished results). This suggests that a high expression is not necessarily related to a maximal rate of translation, and that the choice of colons in artificial illustrative. Although R is rather high (R=38) for this enzyme, which probably leads to the premature termination in titic and in vitro obserred by Manley (1978), a high level of production is obtained (about 3% of total cell proteins). A coexists with premature termination in vivo and in wire at pause sites genes, for example, need not alwars be restricted to the optimal ones

Then Read protein should be considered as a constitutive protein (1000 proteins of E. cali. One hypothesis suggested that it was the plasmid-coded nature the IRNA population of R. coli (Varenne et al., 1982). However, from nucleotide 1983) and B-galactosidase (Kalnins et al., 1983), also have a codon usage quite different from that of proteins like ribosomal proteins (see Table 3 for N). It appears, therefore, that among highly expressed proteins, perhaps constitutive ones should be distinguished from inducible ones with regard to codon usage. copies/cell: Karu & Belk, 1982), although its synthesis can be induced to higher of colicin genes that was responsible for their codon usage not being adapted to equences published recently it can be deduced that chromosomally encoded inducible proteins that are highly expressed, like Pho $oldsymbol{\mathbb{Z}}$ protein (Overbecke of al , Colicin A and E1 are highly expressed proteins after induction (even natural aduction), but have a codon usage different from that of highly expressed

Moreover, these results show that there is no contradiction between discontinuous adapted to this amino acid composition and hads to a fast translation of the non-uniform peptide elongation was clearly demonstrated in vitro and in vivo. The composition of this protein in which Als, Oly and Ser residues account for 87% of message. However, clear parses in translation were observed that probably correspond to the existence of stretches of rare codons intercalated between repetitive domains rich in glyrcine, alanine and serine (Chavaney & Garel, 1981). Little information about discontinuous elongation in cukaryotic cells has been reported. However, in at least two cases, for filtroin (Lizardi et al., 1979, Chavancy & Garel, 1981) and for globin (Protzel & Morris, 1974; Chaney & Morris, 1978), specific case of fibroin is especially interesting because of the peculiar amino seid the amino acid residues. The tRNA population in the posterior silk gland is welland efficient translation.

In conclusion, we presume that also in other prokaryotic and cukaryotic organisms the atochastic search of the ternary complex specific to the codon at the A-site of the ribosome leads to a non-uniform translation. Further experiments will be carried out in our laboratory to confirm this point.

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